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The HPx software for multicomponent reactive transport during variably-saturated flow: Recent developments and applications

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Abstract: HPx is a multicomponent reactive transport model which uses HYDRUS as the flow and transport solver and PHREEQC-3 as the biogeochemical solver. Some recent adaptations have significantly increased the flexibility of the software for different environmental and engineering applications. This paper gives an overview of the most significant changes of HPx, such as coupling transport properties to geochemical state variables, gas diffusion, and transport in two and three dimensions. OpenMP allows for parallel computing using shared memory. Enhancements for scripting may eventually simplify input definitions and create possibilities for defining templates for generic (sub)problems. We included a discussion of root solute uptake and colloid-affected solute transport to show that most or all of the comprehensive features of HYDRUS can be extended with geochemical information. Finally, an example is used to demonstrate how HPx, and similar reactive transport models, can be helpful in implementing different factors relevant for soil organic matter dynamics in soils. HPx offers a unique framework to couple spatial-temporal variations in water contents, temperatures, and water fluxes, with dissolved organic matter and CO₂ transport, as well as bioturbation processes.

Keywords: HYDRUS; HPx; Numerical modeling; Reactive transport; Organic matter; Bioturbation.

INTRODUCTION

Mathematical, process-based simulation models are indispensable tools for elucidating the role of coupled processes in soil systems, including their functioning and providing ecosystem services (Vereecken et al., 2016). Coupled multicomponent reactive transport models for the so-called critical zone (Brantley et al., 2007) are essential tools to analyze nonlinear interactions between hydrological, geochemical and (micro)biological processes and to link the critical zone to other spheres of the earth system such as the hydrosphere, atmosphere, biosphere, and geosphere (Li et al., 2017; Steefel et al., 2005). During the last two decades, a number of tools have been developed to simulate such coupled processes in natural subsurface and engineered systems (Steefel et al., 2015). Numerous applications illustrate the enormous possibilities of these codes (e.g., see references in Li et al. (2017), Steefel et al. (2015)). They typically couple a water flow equation (such as the Richards equation for the vadose zone) with transport equations for heat and solutes based on the advection-dispersion equation, a thermodynamically based geochemical solver, and a kinetic reaction solver, while allowing for the coupling of multiple parameters with a series of state variables (Figure 1). Because of the generic nature of such codes, they provide a flexible (in terms of model formulations) and extendible (in terms of model components) problem-solving environment to tackle important environmental and engineering problems. This paper focuses on the HPx simulator as developed within the HYDRUS computer software packages (Šimůnek et al., 2016). In this introduction we first introduce briefly two aspects which illustrate the need for such modeling tools.

Reactive transport models allow for the direct coupling of time-varying geochemical state variables with transport proper-

ties. In addition to the coupling of microstructural changes due to mineral dissolution and precipitation to porosity and hydraulic properties (Freedman et al. (2004), Wissmeier and Barry (2009), Xie et al. (2015)), fluid properties such as viscosity and density, which are linked to the hydraulic properties of porous media, depend upon the composition of the aqueous solution (Laliberté (2007), Laliberté and Cooper (2004), Sharqawy et al. (2010)). Biomass accumulation or biofilm production during microbiological reactions can also induce changes in hydraulic properties, which may potentially influence the coupled hydrological-physical-geochemical-biological processes operative in porous media and, consequently, such processes as flow paths, the performance of biofilters, and the transport of colloids (e.g., Bozorg et al. (2015a), Bozorg et al. (2015b), Carles Brangari et al. (2017), Or et al. (2007), Rockhold et al. (2002), Yarwood et al. (2006)). The processes of precipitation and dissolution of minerals may be affected by microbiological processes, such as in the engineering practice of microbially induced calcium carbonate precipitation (Hommel et al., 2015; Martinez et al., 2014). Another process that can produce changes in porosity and the hydraulic properties is the migration of fines (solid mineral particles), which is influenced by both hydrodynamic and chemical factors (Bennacer et al., 2017; Mays and Hunt, 2007; Yu et al., 2013). Again, this process is relevant in many engineering applications, but also within the field of soil development. Finally, under saturated conditions, biogenic processes such as methanogenesis and denitrification may result in the formation of gas bubbles, which may result in a reduction of the hydraulic conductivity due to blocking of pore spaces (Amos and Mayer, 2006).

Organic matter dynamics in soils is a crucial process affecting many environmentally-related issues. Degradation produces inorganic carbon, which can be released into the atmosphere.

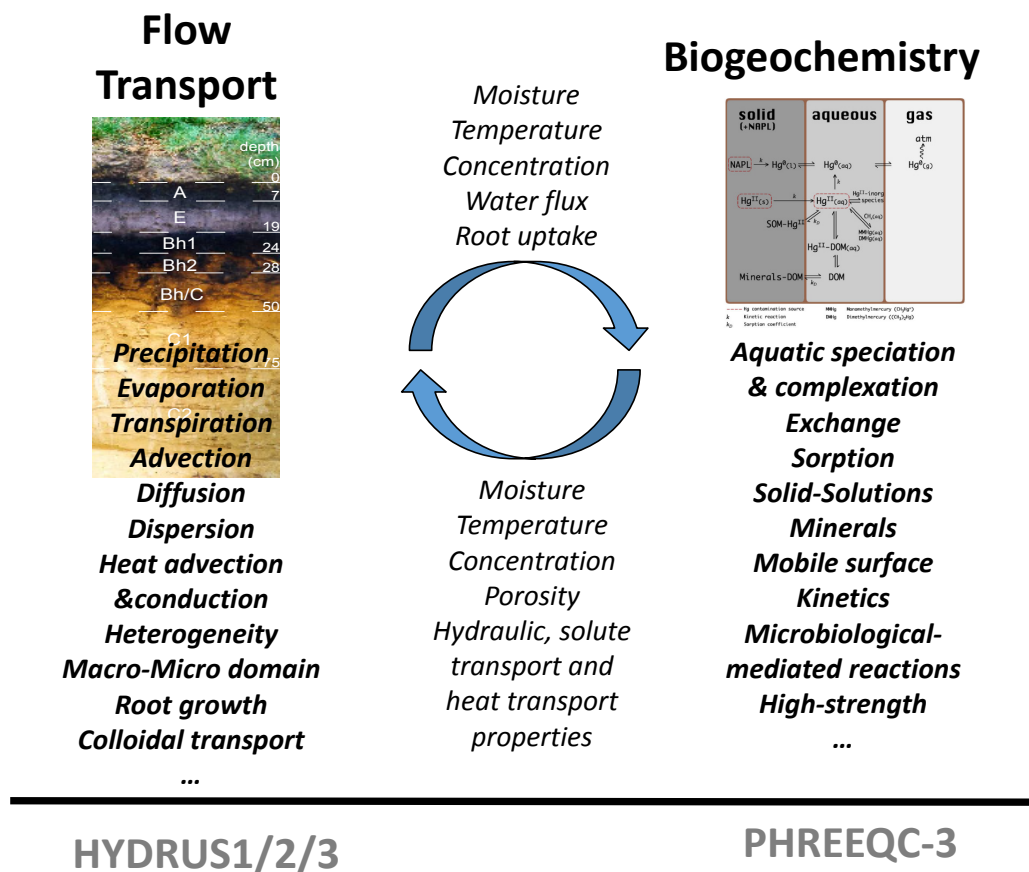


Fig. 1. Reactive transport models coupling water flow and solute transport with biogeochemical processes.

Consequently, CO₂ sequestration in soils may be a factor in controlling atmospheric CO₂, and thus climate change (Todd-Brown et al., 2013). And vice versa, climate change, together with land use, influences soil organic matter (SOM) dynamics (e.g. Batlle-Aguilar et al. (2011)). Dissolved organic matter (DOM) plays also an important role in many soil processes including contaminant transport (e.g., Maes et al. (2006), Paradelo et al. (2012)). For example, Leterme and Jacques (2015) showed that DOM concentrations are among the most sensitive factors in simulating Hg fate in soil systems.

As reported by Manzoni and Porporato (2009), a large number of models exist for simulating SOM dynamics. Many SOM models are so-called compartment models, which divide the heterogeneous nature of SOM into a number of homogeneous pools, and then simulate the fluxes between these pools. Some of these models have a rigid structure in the sense that the number of pools, the abiotic factors that must be considered, their functional forms, and the extent of biotic-abiotic interactions, are fixed. It is therefore not always possible to perform a structured comparison between different types of model formulations. Reactive transport models, on the other hand, provide a flexible structure that enables the introduction of different SOM degradation networks within the same simulation framework. In addition, existing compartment models sometimes strongly simplify flow and transport processes and/or the geochemistry involved, such as simplifying or ignoring pH and buffering effects on degradation or CO₂ geochemistry. For predictions that are more accurate, also within the framework of climate predictions, SOM models should consider temporarily and spatially variable abiotic factors such as temperature and moisture, nutrient dynamics, microbial activity, solid-water interac-

tions, and transport processes of solid and dissolved organic matter and/or inorganic carbon (e.g., Schmidt et al. (2011), Tang and Riley (2015)). Reactive transport models should account for these factors and effects, such as illustrated in recent studies by Tang et al. (2013) and Riley et al. (2014).

The main objective of this paper is to give an overview of new capabilities of the HPx codes compared to the model descriptions given previously by Jacques et al. (2006) and Jacques et al. (2008a). We also wanted to illustrate some of these capabilities as used in several recent studies, as well as present an example involving different SOM degradation networks and bioturbation processes.

THE HPX REACTIVE TRANSPORT CODE

The HPx codes couple the HYDRUS-1D and HYDRUS (2D/3D) software packages (Šimůnek et al., 2016) with PHREEQC (Parkhurst and Appelo, 2013) using a sequential non-iterative coupling approach as explained by Jacques et al. (2006). HYDRUS-1D or HYDRUS (2D/3D) act as the solver for the hydrological and physical processes, including variable-saturated water flow, solute transport, diffusion in the gas phase, and heat transport, whereas PHREEQC is the solver for thermodynamic and kinetic geochemical processes. While the first versions of HPx used PHREEQC-2.4 (Jacques and Šimůnek, 2005; Jacques et al., 2006), the most recent versions 2.4 of HPx use PHREEQC 3.1. Two- and three dimensional flow and transport is included as well (Šimůnek et al., 2012).

Coupling with PHREEQC is performed using so called “hard coupling”, which is in principle similar to the procedures available in the more recently developed wrappers iPhreeqc

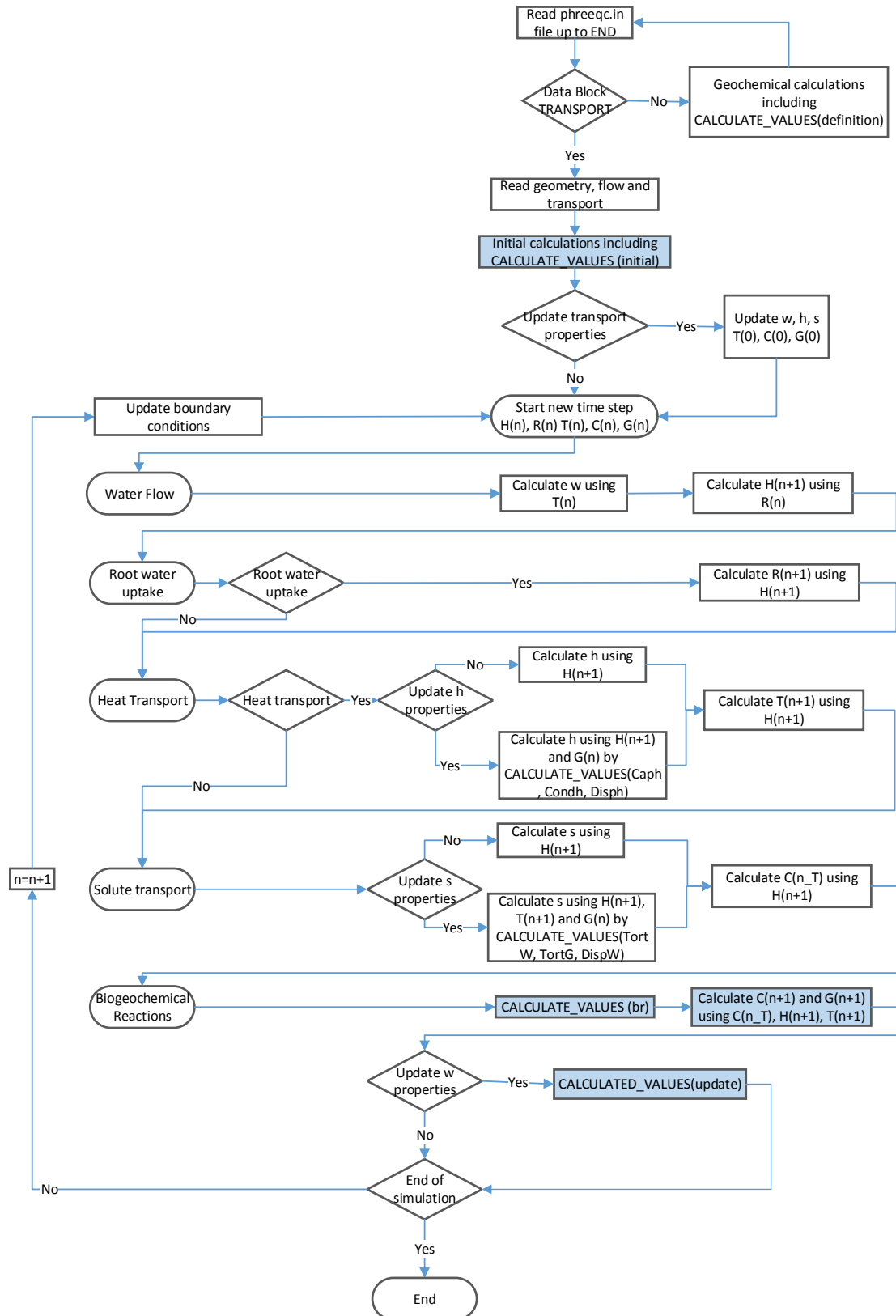


Fig. 2. Flow chart of HPx. The letters w, h, and s indicate parameters for water flow, heat transport, and solute transport, respectively. The variables H, R, T, C, and G indicate state variables for water flow, root water uptake, heat transport, solute transport, and biogeochemical processes, respectively. The boxes with blue background are steps run in parallel using OpenMP. CALCULATED_VALUE(type) indicates the execution of scripts labeled with type.

(Charlton and Parkhurst, 2011) and RMPHREEQC (Parkhurst and Wissmeier, 2015). Due to the hard coupling, HPx does not always include the latest version of PHREEQC. However, HPx includes additional flexibilities compared to the original

PHREEQC, such as linking geochemical definitions (e.g., solution compositions, exchange sites, minerals, kinetic reactants) to the material distribution, or exchanging information between the flow model and the BASIC scripts (e.g., pressure heads,

velocities, and scaling factors). The geochemical input is defined using the standard PHREEQC input conventions using data blocks and identifiers, thereby allowing a direct implementation of the PHREEQC models into HPx. The scripting possibilities are now enhanced by introducing global variables or data tables, which can be used as inline variables or as BASIC scripting variables, using different types of scripting blocks to perform specific calculations, and introducing a specific version of BASIC (such as MyBasic; https://github.com/paladin-t/my_basic) with structured grammar for some specific calculations). Templates for generic types of problems can be generated in advance and then imported into additional input files that can be created during project execution using the INCLUDE\$ statement. Geochemical calculations are performed in parallel using the OpenMP (www.openmp.org) shared memory approach and PHREEQC objects. The main object reads then the input and performs all initial calculations. The information (thermodynamic data and geochemical models) is copied to a user-defined number of additional objects representing chunks in the OpenMP formalism. The nodes are assigned evenly between the chunks. A dynamic schedule controls the calculations of the

nodes in a chunk on the available threads on a computer. After the geochemical calculations, the information from different chunks is assembled and sent to the transport calculations or to the model output files. Because of the OpenMP implementation and the enhanced scripting possibilities, the current flow chart of HPx (see Figure 2) is an extended version of the chart described in Jacques et al. (2006).

Simulation output data now also contain variables, which are integrated over a part of the domain, such as over a particular soil material. HP2 makes it possible to obtain geochemical information along mesh lines or cross-sections at different times as defined in the standard HYDRUS (2D/3D) graphical user interface.

FLOW, TRANSPORT, AND BIOGEOCHEMICAL PROCESSES

Conceptual models

Figure 3 shows schematically the different physical water flow and solute transport models implemented in HPx. The software considers flow, transport, and the prevailing biogeo-

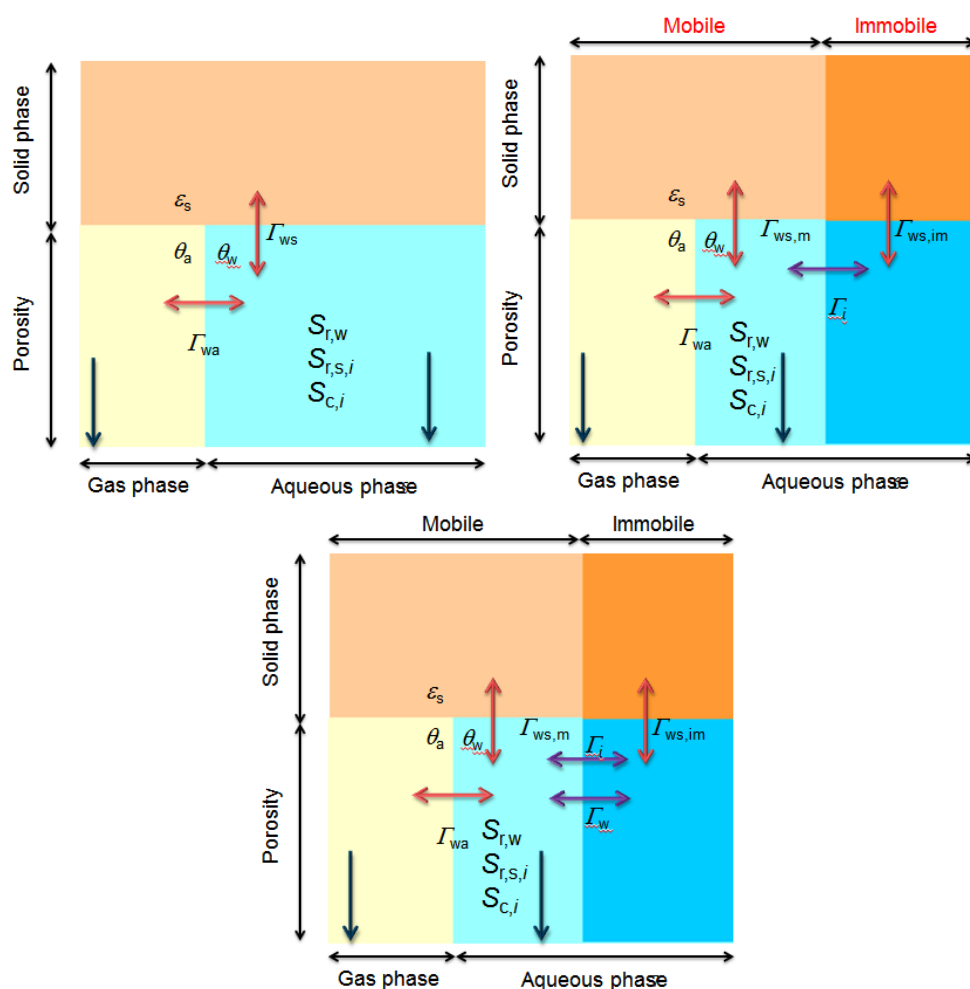


Fig. 3. Conceptual physical and geochemical models implemented in HPx. Black arrows represent transport processes within a single phase, and red arrows exchange processes between different phases. The top left, top right, and bottom plots indicate water flow, solute transport, and biogeochemical reactions in a single-porosity continuum, in a single-porosity continuum with an immobile solute domain, and in a dual-porosity model, respectively (ϵ_s - the solid content [$\text{dm}^3 \text{dm}^{-3}$], θ_a - the air content [$\text{dm}^3 \text{dm}^{-3}$], θ_w - the water content [$\text{dm}^3 \text{dm}^{-3}$], Γ_w - the water exchange between the immobile and mobile domains [$\text{dm}^3 \text{dm}^{-3} \text{T}^{-1}$], Γ_{ws} - the solute exchange between the aqueous phase and the solid phase [$\text{mol dm}^{-3} \text{T}^{-1}$], $\Gamma_{ws,m}$ - the solute exchange between the mobile aqueous phase and the solid phase [$\text{mol dm}^{-3} \text{T}^{-1}$], $\Gamma_{ws,im}$ - the solute exchange between the immobile aqueous phase and the solid phase [$\text{mol dm}^{-3} \text{T}^{-1}$], Γ_i - the solute exchange between the immobile and mobile domains [$\text{mol dm}^{-3} \text{T}^{-1}$], $S_{r,w}$ - root water uptake [$\text{dm}^3 \text{dm}^{-3} \text{T}^{-1}$], $S_{r,s,i}$ - root solute uptake [$\text{mol dm}^{-3} \text{T}^{-1}$], $S_{c,i}$ - kinetic transformations in the aqueous phase [$\text{mol dm}^{-3} \text{T}^{-1}$]).

chemical processes at the continuum scale with three main phases being water (θ_w), air (θ_a) and the solid phase (ϵ_s) as shown in the figure. The water and solid phases in the dual porosity modeling options of the code are divided into two parts, with one part comprising the mobile phase in which water, heat and solutes are allowed to move, and another part being the immobile domain where water is assumed to be stagnant.

Variably-saturated water flow, multicomponent solute transport and heat transport are described using the Richards equation, the advection-dispersion equations (for multiple solutes), and (neglecting energy transported by vapor transport) the convection-dispersion equation, respectively. Transport of reactive gases, which is crucial for simulating biological processes or solid phase alterations such as carbonation of concrete (e.g., Phung et al., 2016) or weathering (Li et al., 2017), is either only by diffusion or advection. The latter is based on the assumptions of negligible air compressibility due to the fast exchange with the atmosphere, zero air flux at the bottom of the profile, and immediate matching of volume changes in the total water content by volume changes in the gas content.

Possible sinks of water in the flow domain are uptake by plants roots ($S_{r,w}$) and exchange between the mobile and immobile water phases (Γ_w), the latter only if dual-porosity water exchange is considered. Sinks or sources of solutes from the aqueous phase are via (i) exchange with roots ($S_{r,i}$ with i referring to a solute component), (ii) kinetic transformations such as radioactive decay, degradation, or redox reactions ($S_{c,i}$), (iii) exchange between the mobile and immobile water phases (Γ_i), and (iv) exchange with the gaseous (Γ_{wa}) or solid phases ($\Gamma_{ws,m}$ and $\Gamma_{ws,im}$) via thermodynamic speciation reactions or kinetic rate equations. The latter reactions represent processes such as sorption, ion exchange, surface complexation, gas-aqueous phase equilibria, and mineral dissolution and precipitation.

Mathematical models

Water flow in its most general form is described using a dual-porosity formulation of the Richards equation (Šimůnek and van Genuchten, 2008):

$$\frac{\partial \theta_m}{\partial t} = \frac{\partial}{\partial x_i} \left[K \left(K_{ij}^A \frac{\partial h}{\partial x_j} + K_{iz}^A \right) \right] - S_{r,w} - \Gamma_w \quad (1)$$

$$\frac{\partial \theta_{im}}{\partial t} = \Gamma_w$$

where θ_m is the volumetric water content in the mobile domain [$L^3 L^{-3}$], θ_{im} is the volumetric water content in the immobile domain [$L^3 L^{-3}$], h is the pressure head [L], x_i ($i = 1, 2, 3$) are the spatial coordinates [L], t is time [T], $S_{r,w}$ is the root water uptake term [T^{-1}], Γ_w is the water exchange term [T^{-1}] driven by a first-order rate equation proportional to the difference in water contents or pressure heads between the mobile and immobile domains, K_{ij}^A are components of a dimensionless anisotropy tensor \mathbf{K}^A , and K is the unsaturated hydraulic conductivity function [LT^{-1}]. The single-porosity models assume that θ_m is equal to the total water content θ , in which case θ_{im} and Γ_m become equal to zero. The second part of Eq. (1) for the immobile region then does not apply.

Several analytical forms are available to describe the unsaturated soil hydraulic properties $\theta(h)$ and $K(h)$ including Brooks and Corey (1964), van Genuchten (1980), Kosugi (1996), and Durner (1994). Root water uptake is driven by potential transpi-

ration. Actual transpiration is calculated based on the spatial-temporal root distribution, while stress factors in the soil are linked to local water contents and salinities. The codes may be used to account for root water uptake compensation by allowing the roots to take up more water in less stressed soil zones (Šimůnek and Hopmans, 2009). Boundary conditions are either system-independent (e.g., constant or variable pressure head (Dirichlet), flux (Neumann), or gradient boundary conditions) or system-dependent by considering atmospheric conditions involving time-dependent precipitation, potential evaporation, and potential transpiration rates.

Heat transport is described as:

$$C_s(\theta) \frac{\partial T}{\partial t} = \frac{\partial}{\partial x_i} \left(\lambda_{ij}(\theta) \frac{\partial T}{\partial x_j} \right) - C_w q_i \frac{\partial T}{\partial x_i} \quad (2)$$

where C_s and C_w are volumetric heat capacities [$ML^{-1}T^{-2}K^{-1}$] of soil and water, respectively, and λ_{ij} is the apparent thermal conductivity of the soil [$MLT^{-3}K^{-1}$]. The reactive module uses temperature to correct equilibrium constants or change rate coefficients as desired using user-defined relations. Equation (2) is valid for both single- and dual-porosity flow models when assuming local temperature equilibrium, i.e., that temperature is the same in the solid phase, as well as in the mobile and immobile liquid phases.

Solute concentrations in the aqueous phase are expressed in terms of total concentrations for a given primary species C_k (mol/kgw), which includes the concentration of all aqueous equilibrium species, c_i (mol/kgw) or so-called secondary species:

$$C_k = c_k + \sum_{l=1}^{N_a} \nu_{lk} c_l \quad (3)$$

where ν_{lk} is the stoichiometric coefficient of species k in secondary species l , $k = 1, \dots, N_c$ (N_c is the number of primary species) and $l = 1, \dots, N_a$ (N_a is the number of secondary species). For each primary species, solute transport in the aqueous phase is described using advection-dispersion-reaction equations of the form:

$$\frac{\partial \theta_m C_{m,k}}{\partial t} = \frac{\partial}{\partial x_i} \left(\theta_m D_{ij}^w \frac{\partial C_{m,k}}{\partial x_j} \right) - \frac{\partial q_i C_k}{\partial x_i} - S_{r,w} C_{r,k} - \Gamma_k - \Gamma_{wa,k} - R_{m,k}^{eq} - R_{m,k}^{kin} \quad (4)$$

$$\frac{\partial \theta_{im} C_{im,k}}{\partial t} = \Gamma_k - R_{im,k}^{eq} - R_{im,k}^{kin}$$

$$\Gamma_k = \omega (C_{m,k} - C_{im,k}) + \Gamma_w (H(\Gamma_w) C_{m,k} + H(-\Gamma_w) C_{im,k})$$

where $C_{m,k}$ and $C_{im,k}$ are the total concentrations of the k species in the mobile and immobile domains [mol/dm^3], respectively, θ_m and θ_{im} have the unit [dm^3/dm^3], D_{ij}^w is the dispersion coefficient tensor [$L^2 T^{-1}$], q_i is the i th component of the Darcian fluid flux density [LT^{-1}], $C_{r,k}$ is the concentration of the sink term [mol/dm^3], Γ_k represents solute exchange between the immobile and mobile domains [$mol/dm^3 T^{-1}$], $\Gamma_{wa,k}$ accounts for the exchange between the aqueous and the gas phase [$mol/dm^3 T^{-1}$], $R_{m,k}^{eq}$, $R_{im,k}^{eq}$, $R_{m,k}^{kin}$ and $R_{im,k}^{kin}$ are aqueous sink/source terms [$mol/dm^3 T^{-1}$], with the superscripts eq and kin referring to equilibrium and kinetic reactions, respectively, and subscripts m and im to the mobile and immobile domains, respectively, ω is the mass transfer coefficient [T^{-1}] and $H()$ is the Heaviside step function.

Reactive gases such as O₂ and CO₂ play important roles in many soil processes (e.g., pyrite oxidation, Mayer et al. (2015)), and hence must form an integral part of reactive transport models. Diffusion in the gas phase is described as:

$$\frac{\partial \theta_a C_{a,k}}{\partial t} = \frac{\partial}{\partial x_i} \left(\theta_a D_{ij}^a \frac{\partial C_{a,k}}{\partial x_j} \right) + \Gamma_{wa,k} \quad (5)$$

where $C_{a,k}$ is the concentration in the gas phase of component k [mol/dm³], D_{ij}^a is the gaseous diffusion tensor [L²T⁻¹], and other symbols as defined before.

The geochemical model calculates aqueous concentrations of the primary and secondary species using an aqueous equilibrium chemical model written as:

$$A_l \rightleftharpoons \sum_{k=1}^{N_c} \nu_{lk} A_k \quad (6)$$

where A_l and A_k are the chemical formulae of the species, ν_{lk} are stoichiometric coefficients and N_c is the number of primary species. Mass action laws permit calculations of the aqueous concentrations as follows:

$$c_l = \frac{\prod_{k=1}^{N_c} (\gamma_k c_k)^{\nu_{lk}}}{K_l \gamma_l} \quad (7)$$

where K_l is the equilibrium constant for the reaction, defined as product of the activities of the reaction products divided by the product of the activities of the reactants, and γ_l and γ_k are the activity correction coefficients to obtain activity a [-] when multiplied with c . The thermodynamic constants are adapted for temperature using the enthalpy of the reaction in the van't Hoff equation or by a six-term analytic equation as a function of temperature (Parkhurst and Appelo (2013), see e.g. also Eq. (2) in the appendix). Some of the parameters of this equation can be linked to thermodynamic variables, such as the change of entropy, enthalpy and heat capacity of reaction, at least for relatively small temperature variations (e.g., Puigdomènech et al. (1997), Jacques (2009)) that are expected for soil conditions. Several activity correction models are implemented, such as ion-association models (Davies equation, extended Debye-Hückel equation), Pitzer model, and a model based on specific ion interaction theory (SIT). Ion exchange, surface complexation, mineral, gas-aqueous phase and solid solutions equilibrium models are available heterogeneous reactions to calculate the $\Gamma_{wa,k}$, $R_{m,k}^{\text{eq}}$ and $R_{\text{im},k}^{\text{eq}}$ terms in Eq. (4).

Equilibrium between the gas and aqueous phases is given as:

$$G_l \rightleftharpoons A_l \quad (8)$$

where G_l is the chemical formula of the gaseous component. The equilibrium constant of the reactions is the ratio of the activity of A_l over the fugacity of G_l . With the assumption of an ideal gas phase, which is valid for low total gas phase pressures, the fugacity coefficient is 1 in which case the fugacity equals the partial pressure of G_l . Fugacity correction factors can be calculated with the Peng and Robinson equation of state (Peng and Robinson, 1976). However, for total gas phase pressures lower than 10 atm, the fugacity coefficient will be close to 1 (Appelo et al., 2014).

A general ion exchange reaction in which the site capacity is always occupied by ions of the opposite charge (i.e., a charge

neutral reaction) is given as:

$$\nu_{X_k} X_k + \nu_l A_l \rightleftharpoons \nu_{X_l} X_l + \nu_k A_k \quad (9)$$

where X denotes the chemical formula of an exchange species and ν is the stoichiometric coefficient, which is linked to a thermodynamic constant using different conventions for the activity correction of the exchange species including Gaines-Thomas, Gapon, and Vanselow convention. The Rothmund-Kornfeld equations (Bloom and Mansell, 2001; Bond, 1995), allows one to consider the effects of a changing selectivity with site occupancy (e.g., Jacques et al. (2012)). Different exchange sites can be combined to define a multisite exchange model (e.g., Jacques et al. (2008b)).

A general equation for surface complexation is:

$$S^{z_s} (A_k^{z_k})_{\nu_k} + \nu_l A_l^{z_l} \rightleftharpoons S^{z_s} (A_l^{z_l})_{\nu_l} + \nu_k A_k^{z_k} \quad (10)$$

where S represents the chemical formula of a surface master species, and z the charge of the surface master species, the aqueous species, or the complexed species. Chemical equilibrium is obtained when the ratio of the activity product is equal to the equilibrium reaction constant, possibly corrected for electrostatic effects. These effects can be either neglected using a non-electrostatic model or incorporated by using a diffuse double layer model (Dzombak and Morel, 1990) or the CD-MUSIC model (Hiemstra and VanRiemsdijk, 1996). The composition of the double layer to counterbalance the surface is estimated by explicitly calculating the charge distribution in the layer (Borkovec and Westall, 1983) or by the Donnan model (Appelo and Wersin, 2007).

A general mineral dissolution/precipitation reaction is of the form:

$$M_l \rightleftharpoons \sum_{k=1}^{N_l} \nu_{kl} A_k \quad (11)$$

where M represents the chemical formula of the mineral and N_l the number of aqueous species in the reaction equation. Since the activity of a pure phase (mineral) is assumed to be equal to 1, the equilibrium constant is the activity product of the aqueous species. In ideal solid solutions, which are homogeneous mixtures of different minerals, the activity of the phase is less than 1, leading to a more stable phase compared to the pure phase. For non-ideal binary solid solutions, described using the Guggenheim series expansion model (Guggenheim, 1937), a mechanical mixture may sometimes be more stable than the homogeneous mixture, resulting in an immiscibility gap or phase separation.

The $R_{m,k}^{\text{kin}}$ and $R_{\text{im},k}^{\text{kin}}$ terms in Eq. (4) represents homogeneous and heterogeneous kinetically controlled reactions, respectively. Homogeneous reaction networks may involve redox, degradation or decay reactions. Each species in such a degradation network must be treated as primary species for which a separate transport equation is required. Typical heterogeneous reactions are dissolution and precipitation of minerals, for which different kinetic expressions exist. One group is based on transition state theory (Aagaard and Helgeson, 1982), which allows for irreversible reactions; the rate parameter may depend on solution composition variables such as pH or activities of other species, which could act as catalyzer or inhibitor. However, irreversible rate equations based on a shrinking core model (e.g., Liu et al. (2014)) or equations in which reaction rates are controlled by microbiological processes can be described as

well. Monod terms for electron acceptors and donors can act as catalytic and inhibitory terms in the microbially-mediated rate equations. Mayer et al. (2002) gave a general rate equation for both surface and transport controlled kinetics for mineral dissolution and precipitation. The scripting possibilities in HPx and PHREEQC permit users to incorporate these or alternative rate equations. In addition, rate equations can be defined for sorption processes when implemented as exchange or surface complexation reactions.

Flow-transport-geochemistry interactions

The concept of scaling hydraulic properties (Vogel et al., 1991), which is typically used in soil sciences to describe their spatial variability, provides a flexible framework for describing the temporal changes of these properties as well. Changes in porosity are then linked to a scaling factor of the water content, α_θ [-], as:

$$\alpha_\theta(t, x) = \frac{\theta_s(t, x) - \theta_r^*}{\theta_s^* - \theta_r^*} \quad (12)$$

where θ_s is the time- and space-variable saturated water content (or the porosity) [$L^3 L^{-3}$], and θ_s^* [$L^3 L^{-3}$] and θ_r^* [$L^3 L^{-3}$] are the reference saturated and residual water contents, respectively, for a specific material. Variations in the hydraulic conductivity in time and space, $K(x, t)$ [$L T^{-1}$], are described by a scaling factor of the hydraulic conductivity, α_K [-], as follows

$$\alpha_K(t, x) = \frac{K_s(t, x)}{K_s^*} \quad (13)$$

where K_s^* is the reference saturated hydraulic conductivity for a specific material [$L T^{-1}$]. This formulation enables one to include the effects of temperature and geochemistry on viscosity and density, as well as of precipitation/dissolution/clogging processes on the microstructure. Also, changes in the apparent air-entry value can be described using a scaling factor for the pressure head, α_h [-], such that h/α_h rather than h (for the reference hydraulic properties) is used to obtain the water content and hydraulic conductivity. HPx does not provide any functional relation for calculating the scaling factors. Via the input file, users can implement their preferred model relating geochemical variables to the scaling factors. In the current implementation, change of porosity and scaling factor for the pressure head (in the dual porosity model with water exchange driven by pressure head gradient) are limited to the mobile domain. Assuming a constant porosity of the immobile domain, the change in α_h reflects a change in porosity of the mobile domain only. One limitation is that the scaling factors only describe linear changes in the hydraulic properties. For example, they do not allow for capturing a model for hydraulic property changes which assumes that precipitation and dissolution reactions occur only in the fully water-saturated part of the pore space (e.g., the model of Wissmeier and Barry (2009)). However, when the reactions take place in all of the pore space, the linear scaling model is applicable (Wissmeier and Barry, 2010).

HPx can also update solute and heat transport properties such as the tortuosity of the aqueous and gaseous phases, and the dispersivity, heat capacity, heat conductivity, and heat dispersivity. Tortuosity is often linked to porosity using Archie's law or the model of Millington and Quirk (1961). For cement systems, some models discriminate between the types of porosity (gel versus capillary porosity) and associated cement

phases (Patel et al., 2016). The heat capacity can be linked to individual heat capacities of the minerals in the system. However, not much literature evidence exists for a model linking dispersivity and conductivity to geochemical state variables.

Root solute uptake

Kinetic uptake and exudation processes can be implemented in HPx using rate expressions defined in the input files. Examples of PHREEQC implementations of these types of processes were given by Seuntjens et al. (2004) and Nowack et al. (2006). Compared to the HYDRUS implementations, also available in HPx, some additional features can be taken into account when the rate equations $R_{n,k}^{\text{kin}}$ in Eq. (4) are used, such as species-specific uptake rather than uptake of an element irrespective of the distribution among different aqueous species, inclusion of counter-ions, root exudation, and uptake parameters depending upon soil geochemistry. It is also possible to incorporate the term $S_{r,w}C_{r,k}$ in Eq. (4) in $R_{n,k}^{\text{kin}}$. Following Silberbush et al. (2005), a general formulation of root solute uptake that combines passive and active uptake is given by

$$r_l = p_l + a_l \quad (14)$$

$$= p_l + A_{\text{root}} J_{l,\text{max}} \left(\omega_{\text{gc}} \right) \frac{c_l - c_{l,\text{min}}}{K_{l,m} \left(\omega_{\text{gc}} \right) + (c_l - c_{l,\text{min}})}$$

$$p_{a,l} = \min[S_{w,r} \min[c_l, c_{\text{max},l}], \quad (15)$$

$$A_{\text{root}} J_{p,\text{max},l} \left(\omega_{\text{gc}} \right),$$

$$p_{p,l}] \quad \text{if } c_l > c_{r,l}$$

$$= 0 \quad \text{otherwise}$$

where r_l refers to the uptake of a species l , p_l and a_l are passive and active uptake rates [$\text{mol}/\text{dm}^3 T^{-1}$], $c_{\text{max},l}$ is the maximum allowed concentration for passive uptake [mol/dm^3], A_{root} is the root surface area [L^2/dm^3], $J_{p,\text{max},l}$ is the maximum allowed passive nutrient uptake rate [$\text{mol } L^{-2} T^{-1}$], ω is the current geochemical condition, $p_{p,l}$ is the potential passive nutrient uptake rate [$\text{mol}/\text{dm}^3 T^{-1}$], $c_{r,l}$ is the critical concentration below which passive root uptake is zero [mol/dm^3], $J_{l,\text{max}}$ is the maximum active nutrient uptake which may depend on geochemical conditions or the maximum potential active uptake [T^{-1}], $K_{l,m}$ is the Michaelis-Menten parameter [mol/dm^3], and $c_{l,\text{min}}$ is the minimum nutrient concentration below which active uptake is zero. The expression for passive uptake is generic as it limits uptake by (i) a maximum allowable concentration for uptake, (ii) a maximum nutrient uptake flux, and (iii) the maximum potential root uptake rate. It is important to note that it is not straightforward to implement active and passive uptake processes as discussed by Šimůnek and Hopmans (2009), in which active uptake is only invoked when passive uptake does not meet a maximum nutrient plant demand.

OVERVIEW OF RECENT APPLICATIONS

The early studies of Jacques et al. (2008a, 2008b) illustrated the coupled effects of water flow dynamics and geochemistry on the mobility and availability of various contaminants. Since water content variations due to the temporal nature of precipitation, infiltration, and evapotranspiration continually change concentrations in the aqueous phase, other chemical factors and processes such as ionic strength, pH, and aqueous complexation are affected as well. Differences in component mobility result in changes in aqueous speciation and, consequently, both

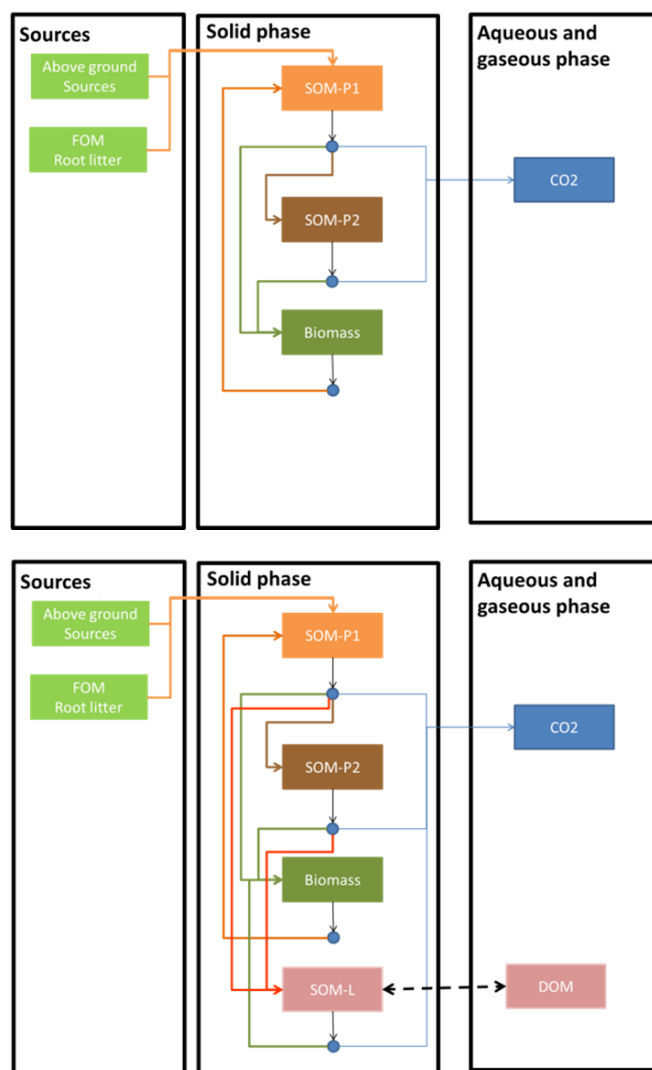


Fig. 4. Organic matter decomposition networks: (top) a scheme without leachable soil organic matter, (bottom) a scheme with leachable soil organic matter. Orange lines indicate decomposition to easily decomposable soil organic matter (SOM-P1), brown lines indicate decomposition to slowly decomposable soil organic matter (SOM-P2), green lines indicate assimilation into biomass (Biomass), blue lines indicate respiration and production of inorganic carbon species (CO_2), and red lines indicate decomposition towards leachable soil organic matter (SOM-L). The dashed black line represents kinetic exchange between leachable soil organic matter and dissolved organic matter.

mineral dissolution/precipitation and sorption can be affected by the temporal-spatial variability of water contents and water fluxes. Comprehensive studies and/or assessments of fate and transport of contaminants in soil systems need to take into account these types of variations. HPx already has been applied in many studies, including for evaluation of Hg fate in soil systems (Bessinger and Marks, 2010; Leterme and Jacques, 2015; Leterme et al., 2014) or, more recently, scenario analyses involving the transport of trace metals in a soil leached with coal seam gas produced water (Mallants et al., 2017). The benchmark study of Greskowiak et al. (2015), in which HPx was used as one of several reactive transport models they evaluated, illustrated the applicability of HPx to describe U transport using a multi-rate mass transfer model (MRMT, e.g., Haggerty and Gorelick (1995)).

Another benchmark study was used to verify the implementation of changes in the porosity, diffusion coefficient and hydraulic conductivity as a consequence of mineral precipitation and dissolution in multiple reactive transport models (Xie et al., 2015), including HP1. Valdes-Abellan et al. (2017) developed a model (using HP1) to simulate the spatial-temporal evolution of soil hydraulic properties during non-conventional water irrigation during non-isothermal transient flow. In addition to linking the porosity, α_h , and the saturated hydraulic conductivity to the amount of precipitated or dissolved minerals, soil hydraulic properties were also linked to changes in the soil water composition, salinity, and temperature. Valdes-Abellan et al. (2017) in their study indicated that a close interplay exist between gypsum-calcite dissolution/precipitation, evaporation fronts, and the key role of the partial pressure of CO_2 . While the partial pressure of CO_2 was linked only to calcite equilibria, the role of organic matter degradation (and corresponding CO_2 production) could potentially be incorporated within the HPx framework as well (see below).

HYDRUS in its specialized C-Ride add-on module solves a set of equations describing the transport of colloid and colloid-facilitated solute transport (Šimůnek et al., 2006). A similar set of equations can be implemented in HPx, thus allowing one to consider the effects of geochemistry on various parameters such as attachment efficiency or competitive sorption on the transport of colloids. Zhou et al. (2016) developed a model simulating the transport of three antibiotics in the presence of different kinds of natural organics originating from manure. Compared to the comprehensive capabilities in the standard HYDRUS code, HPx can account for competitive sorption of antibiotics to the soil using a Langmuir competitive kinetic adsorption model in conjunction with competitive kinetic sorption on mobile and immobile organic colloids. The role of geochemistry on colloid mobility was described in a study by Makselon et al. (2017) who modeled the transport of silver nanoparticles in an experiment with flow interruption, evaporation, and ionic strength changes. Ionic strength, and hence the general aqueous geochemical conditions, determined the value of the attachment efficiency, which can be calculated using DVLO theory. A lower ionic strength results in a lower attachment efficiency because of a higher energy barrier and a smaller secondary minimum. BASIC scripts in the input file included root finding algorithms to identify the primary and secondary minima to obtain the attachment efficiency as a function of the changing geochemical conditions during the transport experiment.

EXAMPLES

Organic matter dynamics

As an illustration of the flexible framework of HPx to define reaction networks in general, and for soil organic matter (SOM) degradation in particular, a reaction network loosely based on a study by Porporato et al. (2003) was implemented for this example (HPx projects are available at <https://www.pc-progress.com/en/Default.aspx?h1d-library>). For convenience, the single-porosity model formulation is used in this example. Figure 4 shows the network with three organic matter pools: (i) a fast decomposable SOM pool (SOM-P1) with concentration $C_{\text{SOM-P1}}$, (ii) a slow decomposable SOM pool (SOM-P2) with concentration $C_{\text{SOM-P2}}$, and (iii) a biomass pool (Biomass) with concentration C_B . In addition, inorganic carbon is released in the aqueous and gaseous soil phases during decomposition, C_i .

This decomposition is described in terms of four ordinary differential equations:

$$\frac{dC_{\text{SOM-P1}}}{dt} = i_F + d_B - d_{\text{SOM-P1}} \quad (16)$$

$$\frac{dC_{\text{SOM-P2}}}{dt} = r_{\text{SOM-P1} \rightarrow \text{P2}} d_{\text{SOM-P1}} - d_{\text{SOM-P2}} \quad (17)$$

$$\frac{dC_B}{dt} = r_{\text{SOM-P1} \rightarrow B} d_{\text{SOM-P1}} + r_{\text{SOM-P2} \rightarrow B} d_{\text{SOM-P2}} - d_B \quad (18)$$

$$R_{m,C(4)}^{\text{kin}} = r_{\text{SOM-P1} \rightarrow I} d_{\text{SOM-P1}} + r_{\text{SOM-P2} \rightarrow I} d_{\text{SOM-P2}} \quad (19)$$

where d_X refers to the decomposition rates, $r_{x \rightarrow y}$ denotes the fraction of the decomposition products of pool x going to pool y , and i_F denotes the addition of fresh organic matter as litter fall on top of the soil profile or as root decay. Wutzler and Reichstein (2008) discussed different model formulations that can be used to include decomposer biomass into the decomposition rates: (i) no representation of the biomass using equations that are based only on the substrate, (ii) a linear representation of biomass, and (iii) a nonlinear representation of biomass. These models differ in the scaling behavior with the substrate and how they represent priming effects. Although all of these formulations could be included in HPx, the example given below is limited to a non-explicit representation of the biomass and a linear model of the available substrate, i.e.,

$$d_X = k_X C_X \quad (20)$$

where k_X is the first-order decomposition term [T^{-1}], C_X is the concentration of the substrate [ML^{-3}], and X denotes a soil organic matter pool. The value of the rate parameter k_X may depend on temperature $f_T(T)$ [-] and soil moisture $f_M(S_\theta)$ [-] as follows:

$$k_X = k_{X,0} f_T(T) f_M(S_\theta) \quad (21)$$

where $k_{X,0}$ is the reference rate [T^{-1}]. Sierra et al. (2012) gave an overview of different functions to account for the effects of environmental factors on the degradation rate. Here we describe $f_M(S_\theta)$ using the equations of Porporato et al. (2003):

$$f_M(S_\theta) = \begin{cases} \frac{S_\theta}{S_{fc}} & S_\theta \leq S_{fc} \\ \frac{S_{fc}}{S_\theta} & S_\theta > S_{fc} \end{cases} \quad (22)$$

where S_θ is the degree of saturation ($=\theta/\theta_s$) [-] and S_{fc} is the saturation degree at field capacity [-]. The temperature dependency is described using the formulation from ROTC (Jenkinson et al., 1990):

$$f_T(T) = \frac{47.9}{1 + \exp\left(\frac{106}{T + 18.3}\right)} \quad (23)$$

Still needed to complete the description of the organic matter decomposition network is an aqueous speciation model for inorganic C species. We refer to the Appendix for details. Table 2 further gives the parameters used in a transient flow simulation for a meteorological time series representative of Belgium conditions.

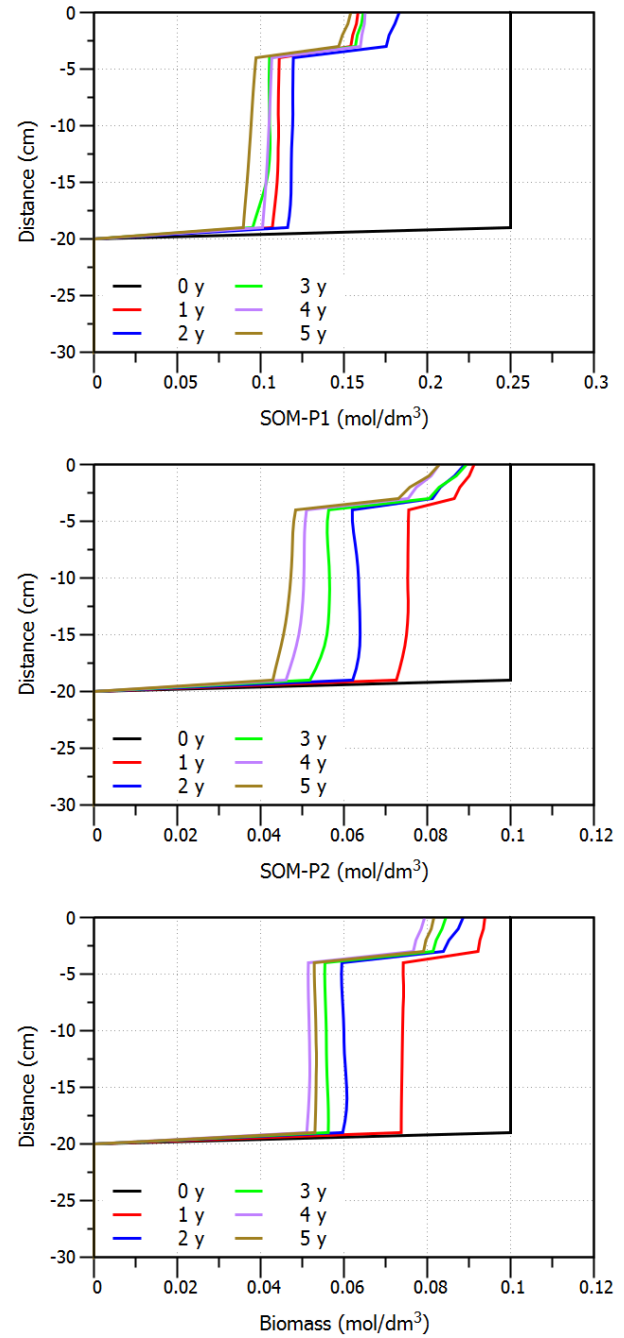


Fig. 5. Profiles of SOM-P1 (top), SOM-P2 (middle), and Biomass (bottom) for network 1.

Figure 5 shows the computed distributions versus depth of the various pools of soil organic matter and biomass during five consecutive years. Large changes during the first few years indicate that selected initial and production terms are not in long-term equilibrium with the degradation processes. The current set of parameters and meteorological variables do not produce much variability in the vertical direction. The relative enrichment of the top 5 cm is due to SOM-P1 originating both from litter fall and root decay whereas deeper in the profile, only root decay contributes to the SOM-P1 pool. A more thorough analysis of the model setup using a sensitivity analysis of the model parameters is beyond the scope of the current paper but could be done straightforwardly with HPx as was illustrated by Leterme and Jacques (2015).

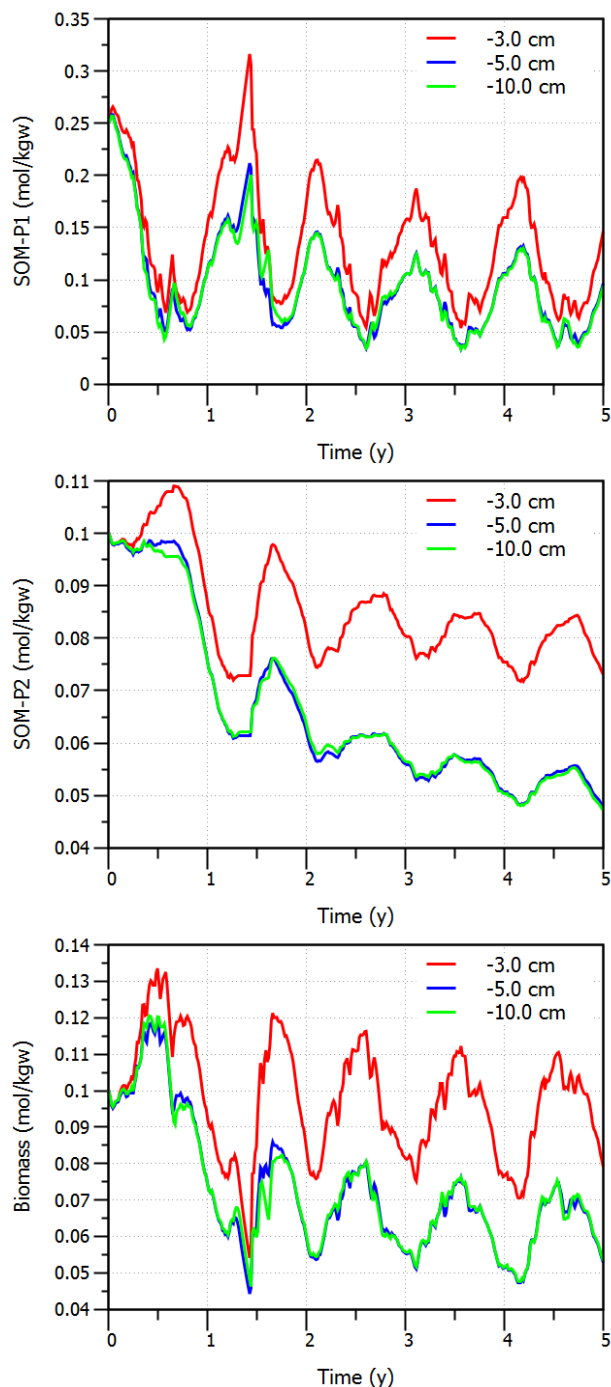


Fig. 6. Time series of SOM-P1 (top), SOM-P2 (middle), and Biomass (bottom) for network 1.

Figure 6 shows time series of various pools of SOM and Biomass at several depths. The strong seasonal fluctuations in these pools are due to variations in soil water contents and temperatures, leading to changes in the environmental factors on the degradation rates as expressed by Eqs. (21)–(23). As shown in Figure 7, variations in f_M and f_T are similar at different soil depths, which explain the relatively uniform distribution of various organic pools (Figure 5). HPx in general provides geochemical information of the system such as pH (not further shown here) and inorganic aqueous and gaseous C concentrations, the latter shown in Figure 8 in terms of the partial pressure of CO_2 in the soil air phase. In this example C was the only aqueous species and no pH controlling

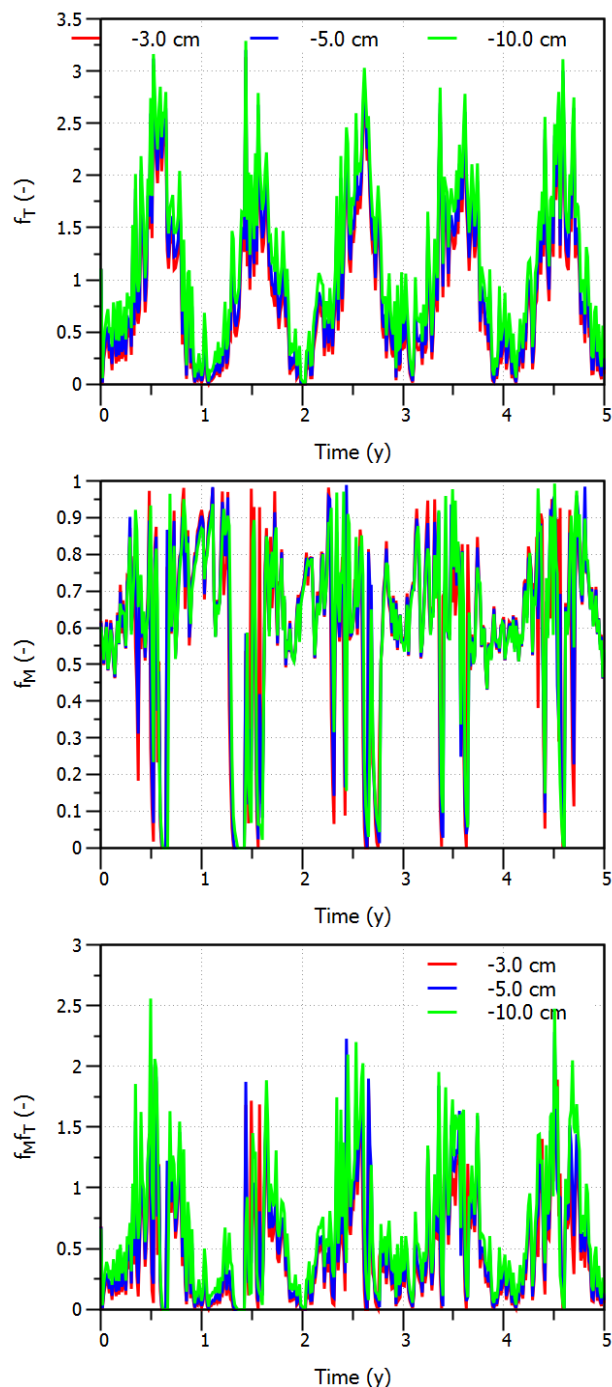


Fig. 7. Time series of f_M (top), f_T (middle), and $f_M f_T$ (bottom) for network 1.

components or processes were included. Figure 8 also shows the cumulative flux of CO_2 from the soil to the atmosphere, which is a critical variable in soil-plant-atmosphere and regional or global climate studies (e.g., Thaysen et al. (2014)).

Dissolved organic matter

The previous organic matter degradation model is further extended in this example by including a leachable organic matter pool (SOM-L) and a dissolved organic matter pool (DOM), which are both kinetically exchanged. The formulation, loosely based on the model of Braakhkke et al. (2011), allows for

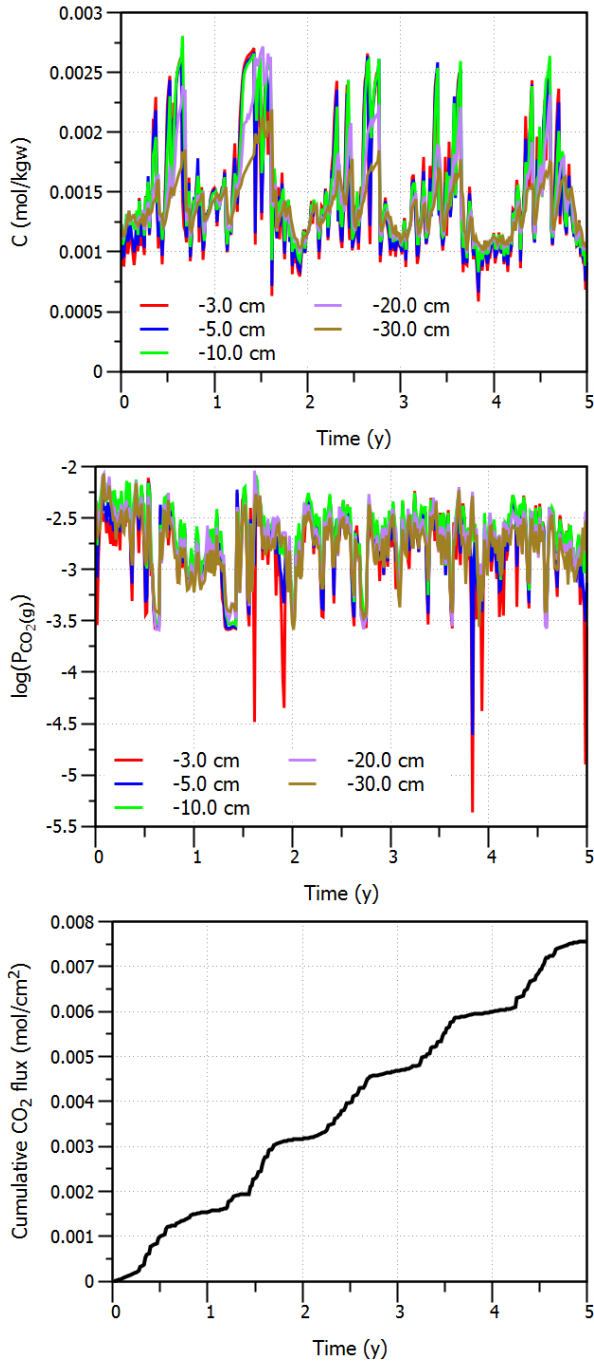


Fig. 8. Time series of C concentration (mol/kgw) (top), P_{CO_2} (middle) and cumulative CO_2 flux to the atmosphere (bottom) for network 1.

advective-dispersive transport of DOM. The set of governing ordinary-differential equations is as follows:

$$\frac{dC_{SOM-P1}}{dt} = i_F + d_B - d_{SOM-P1} \quad (24)$$

$$\frac{dC_{SOM-P2}}{dt} = r_{SOM-P1 \rightarrow P2} d_{SOM-P1} - d_{SOM-P2} \quad (25)$$

$$\begin{aligned} \frac{dC_B}{dt} = & r_{SOM-P1 \rightarrow B} d_{SOM-P1} + r_{SOM-P2 \rightarrow B} d_{SOM-P2} \\ & + r_{SOM-L \rightarrow B} d_{SOM-L} - d_B \end{aligned} \quad (26)$$

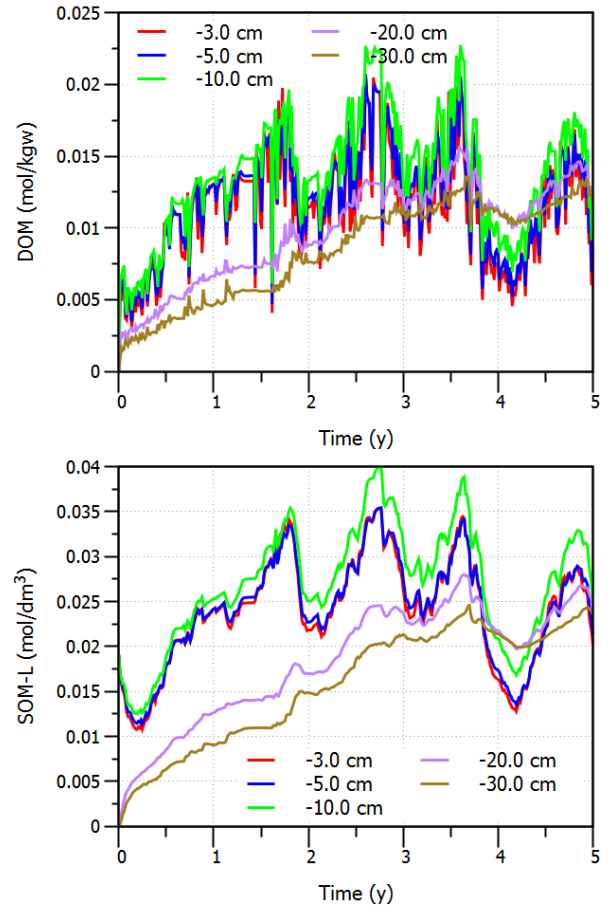


Fig. 9. Time series of DOM (top) and SOM-L (bottom) for network 2.

$$\begin{aligned} \frac{dC_{SOM-L}}{dt} = & r_{SOM-P1 \rightarrow SOM-L} d_{SOM-P1} + r_{SOM-P2 \rightarrow SOM-L} d_{SOM-P2} \\ & - d_{SOM-L} - d_S \end{aligned} \quad (27)$$

$$R_{m,C(4)}^{kin} = r_{SOM-P1 \rightarrow I} d_{SOM-P1} + r_{SOM-P2 \rightarrow I} d_{SOM-P2} + r_{SOM-L \rightarrow I} d_{SOM-L} \quad (28)$$

$$R_{m,DOM}^{kin} = d_S \quad (29)$$

where d_S is the desorption term:

$$\begin{aligned} d_S = & k_S (C_{SOM-L} - C_{SOM-L}^{eq}) \\ = & k_S \left(C_{SOM-L} - \frac{K_L S_{max} C_{DOM}}{1 + K_L C_{DOM}} \right) \end{aligned} \quad (30)$$

in which K_L is the coefficient of binding affinity [$dm^3 gC^{-1}$] and S_{max} the maximum adsorption capacity [$gC dm^{-3}$]. Time series of the new two pools, SOM-L and DOM, are shown in Figure 9. DOM shows short-scale temporal variations due to changes in the water content. These variations are less pronounced for SOM-L due to the kinetics of the sorption process. The advection and dispersion of DOM result in an increase in both DOM and SOM-L at depths below the initial source of soil organic matter between 0 and 20 cm below the soil surface. Note that the other pools potentially will develop also below the 20-cm depth when this degradation scheme is used, but their amounts remain very small for the chosen set of parameters.

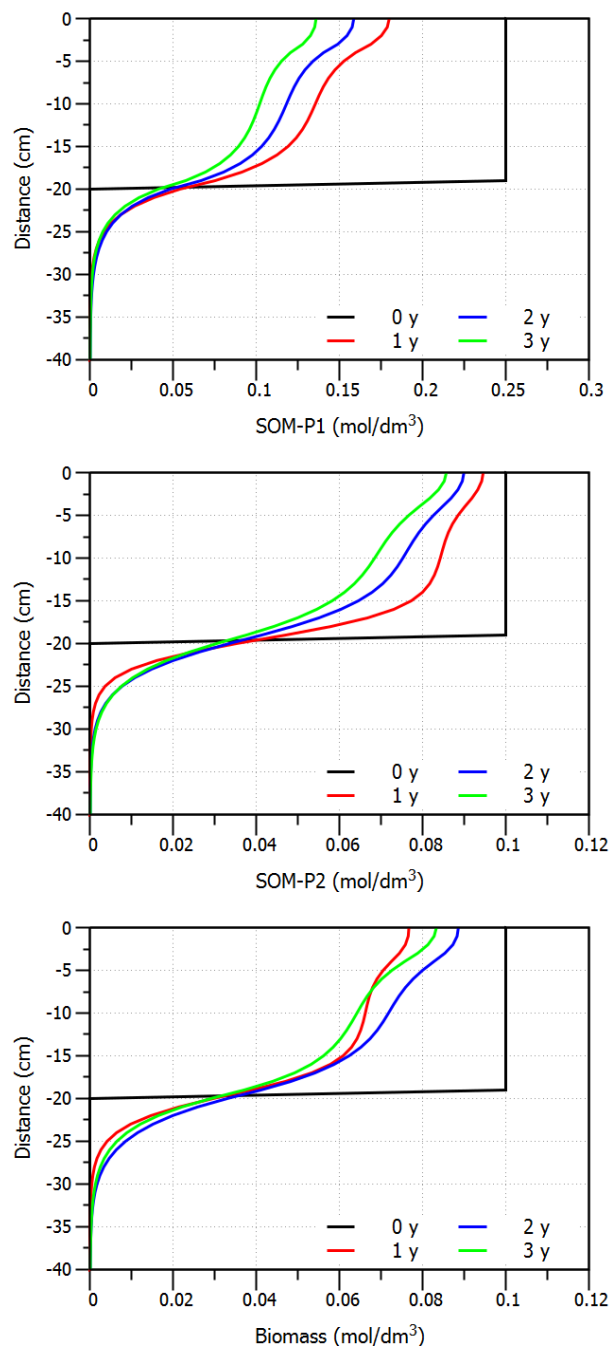


Fig. 10. Profiles of SOM-S1 (top), SOM-S2 (middle) and Biomass (bottom) for a simulation with network 1 including biodiffusion.

Bioturbation

The long term prediction of soil carbon cycling requires an explicit representation of the vertical distribution of soil organic matter, and hence a model able to represent processes leading to the vertical distribution of SOM such as the root litter distribution and the transport of soil organic matter (Braakhekke et al., 2011). The biological transport of soil solid phases, often referred to as bioturbation, depends strongly on the behavior of the biological community species, which necessitated the development of different modeling approaches (Meysman et al., 2003a). Models based on Fick's law are local random models (Meysman et al., 2005), whereas non-local models need to be employed when ingestion and egestion processes of SOM by bio fauna are vertically separated (Jarvis et al., 2010).

As an example, a biodiffusion model was implemented within the input files of HPx:

$$J_{s,X} = -D_{B,X} \frac{\partial C_X}{\partial x} \quad (31)$$

where $J_{s,X}$ is the solid phase diffusive flux of the X th pool [$\text{mol T}^{-1} \text{L}^{-2}$], $D_{B,X}$ is the effective solid phase diffusion coefficient [$\text{L}^2 \text{T}^{-1}$], and C_X is the solid phase concentration [mol L^{-3}]. The biodiffusion model was applied to organic network 1 described above with SOM-P1, SOM-P2, and biomass. We assumed rather arbitrarily that $D_{B,X} = 10^{-10} \text{ m}^2/\text{s}$. As shown in Figure 10, a smooth profile of the three organic pools developed over the course of 3 years to about 15 cm below the initial layer with organic matter.

CONCLUSIONS

The original HP1 model (Jacques and Šimůnek, 2005; Jacques et al., 2006) has seen many extensions during the past 10 years or more. The most important changes were the development of two- and three-dimensional versions (HPx), and the incorporation of the software PHREEQC-3, OpenMP, enhanced scripting possibilities, diffusion in the gas phase of reactive gases in the software, and linking geochemical state variables to transport properties. Using the terminology of Meysman et al. (2003b), HPx is now moving towards a more flexible and extendible problem-solving environment to allow users to incorporate a range of user-defined conceptual models to tackle their specific projects. Flexibility and extendibility are discussed in this manuscript with respect to coupling geochemistry with transport properties, colloid and colloid-affected transport, bioturbation, and root solute uptake. Recent applications show that HPx, and similar reactive transport codes, are useful tools to evaluate many environmental and engineering problems both from a scientific as well as from an application point of view, thus underpinning the main statement from Li et al. (2017).

In this paper, we illustrated the capabilities and flexibility of HPx by developing a reaction-transport model for soil organic matter, in which we included different factors important in soil organic matter degradation. Only a few SOM degradation models can similarly consider spatial-temporal variations in water contents, water fluxes, and temperatures, as well as different conceptual models for SOM degradation networks, advection-dispersive transport of DOM, CO_2 transport, and bioturbation in one modeling framework. Within the scope of this paper, the role of microbial processes together with the transport of oxygen were not addressed, but conceptually could be implemented in a very similar way. We also acknowledge that further parameter sensitivity studies and validation of these models are crucial in further model applications.

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APPENDIX – Details about the organic matter degradation model.

The aquatic geochemical reactions within the adopted modeling approach are limited to the inorganic carbon reactions, as given in Table 1. For all aqueous species, the activity correction model of WATEQ is used:

$$\log \gamma = \frac{-Az^2\sqrt{\mu}}{1+Bz\sqrt{\mu}} + b\mu \quad (1)$$

where γ is the activity correction coefficient, z is the charge, μ is the ionic strength, A and B are constants at a given temperature, and a and b are species-dependent parameters, except for CO_2 for which the activity coefficient is obtained as $\log \gamma = 0.1 \mu$. The temperature dependence of the mass action constant is given as:

$$\log_{10} K = A_1 + A_2T + \frac{A_3}{T} + A_4 \log_{10} T + \frac{A_5}{T^2} + A_6T^2 \quad (2)$$

where A_1, \dots, A_6 are parameters and T is temperature in Kelvin. The relevant temperature range is limited between ~ 0 to $\sim 30^\circ\text{C}$.

Tables 2 through 5 provide a list of parameter definitions and values used in the simulations of the organic matter degradation networks.

Table 1. Overview of species, geochemical reactions, and thermodynamic parameters (values are from the phreeqc.dat database as incorporated in PHREEQC-3).

Species	Reaction	Activity correction		Temperature dependence					
		<i>a</i>	<i>b</i>	<i>A</i> ₁	<i>A</i> ₂	<i>A</i> ₃	<i>A</i> ₄	<i>A</i> ₅	<i>A</i> ₆
H ⁺		9.0	0						
CO ₃ ²⁻		5.4	0						
OH ⁻	H ₂ O = OH ⁻ + H ⁺	3.5	0	293.29	0.1361	-10576.9	-123.73	0	-6.996e-5
HCO ₃ ⁻	CO ₃ ²⁻ + H ⁺ = HCO ₃ ⁻	5.4	0	107.89	0.0325	-5151.8	-38.93	563713.9	
CO ₂	CO ₃ ²⁻ + 2H ⁺ = CO ₂ + H ₂ O	–	–	464.20	0.0934	-26986.2	-165.76	2248628.9	
CO _{2(g)}	CO ₂ = CO ₂	–	–	119.87	2.19e-2	-7337.8	-44.77	669371	

Table 2. Parameter values for network 1 (Eqs. (16)–(19)). Parameter values are for illustrative purpose.

Symbol	Parameter description	Value	Unit
<i>i</i> _F	Addition of fresh organic matter	0.0015	gC/d/dm ³ bulk
<i>k</i> _B	Decay rate for biomass pool	8.5×10 ⁻³	1/d
<i>r</i> _{SOM-P1→P2}	Transformation factor from SOM-P1 to SOM-P2	0.1	–
<i>r</i> _{SOM-P1→B}	Transformation factor from SOM-P1 to B	0.3	–
<i>r</i> _{SOM-P1→I}	Transformation factor from SOM-P1 to I	0.6	–
<i>r</i> _{SOM-P2→B}	Transformation factor from SOM-P2 to B	0.4	–
<i>r</i> _{SOM-P2→I}	Transformation factor from SOM-P2 to I	0.6	–
<i>ρ</i> _b	Bulk density	1.43	kg/dm ³ bulk
<i>k</i> _{SOM-P1}	Rate constant for SOM-P1 degradation	3.00×10 ⁻²	1/d
<i>k</i> _{SOM-P2}	Rate constant for SOM-P2 degradation	5.00×10 ⁻³	1/d

Table 3. Parameter values for network 2 (Eqs. (24)–(30)). Parameter values are for illustrative purpose.

Symbol	Parameter description	Value	Unit
<i>i</i> _F	Addition of fresh organic matter	0.0015	gC/d/dm ³ bulk
<i>k</i> _B	Decay rate for biomass pool	8.5×10 ⁻³	gC/d/dm ³ bulk
<i>r</i> _{SOM-P1→P2}	Transformation factor from SOM-P1 to SOM-P2	0.1	–
<i>r</i> _{SOM-P1→B}	Transformation factor from SOM-P1 to B	0.3	–
<i>r</i> _{SOM-P1→I}	Transformation factor from SOM-P1 to I	0.55	–
<i>r</i> _{SOM-P1→L}	Transformation factor from SOM-P1 to SOM-L	0.05	–
<i>r</i> _{SOM-P2→B}	Transformation factor from SOM-P2 to B	0.4	–
<i>r</i> _{SOM-P2→I}	Transformation factor from SOM-P2 to I	0.5	–
<i>r</i> _{SOM-P2→L}	Transformation factor from SOM-P2 to SOM-L	0.1	–
<i>r</i> _{SOM-L→B}	Transformation factor from SOM-P2 to B	0.1	–
<i>r</i> _{SOM-L→I}	Transformation factor from SOM-P2 to I	0.9	–
<i>K</i> _L	Binding affinity of DOM	5	dm ³ /gC
<i>S</i> _{max}	Binding capacity for DOM	0.4	gC/dm ³ bulk
<i>ρ</i> _b	Bulk density	1.43	kg/dm ³
<i>k</i> _{sor}	First-order kinetic sorption term	0.05	d ⁻¹
<i>k</i> _{SOM-P1}	Rate constant for SOM-P1 degradation	3.0×10 ⁻²	1/d
<i>k</i> _{SOM-P2}	Rate constant for SOM-P2 degradation	5.0×10 ⁻³	1/d
<i>k</i> _{SOM-L}	Rate constant for SOM-L degradation	2.0×10 ⁻⁶	1/d

Table 4. Initial organic matter pools.

Organic Pool	Value	Unit
SOM-P1	1.75×10 ⁻⁴	kgC/kg soil
SOM-P2	7.00×10 ⁻⁵	kgC/kg soil
SOM-L1 [§]	0.019	gC/dm ³ bulk
B	7.00×10 ⁻⁵	kgC / kg soil
DOM [§]	0.01	gC/dm ³ soil

[§] only for reaction network 2 * Calculated in equilibrium with DOM

Table 5. Soil hydraulic parameters for the van Genuchten (1980) soil hydraulic functions, solute transport parameters, and plant uptake parameters (for the S-shape water uptake reduction model – see the HYDRUS-1D manual for more information Šimunek et al. (2013)) for single porosity model implementation.

Symbol	Parameter description	Value	Unit
<i>θ</i> _r	Residual water content	0.08	dm ³ /dm ³
<i>θ</i> _s	Saturated water content	0.45	dm ³ /dm ³
<i>α</i>	Shape parameter of the water retention function	0.05	cm ⁻¹
<i>n</i>	Shape parameter of the water retention functions	1.6	–
<i>K</i> _s	Saturated hydraulic conductivity	25	cm day ⁻¹
<i>D</i> ^w	Diffusion coefficient in the aqueous phase	2	cm ² day ⁻¹
<i>D</i> ^a	Diffusion coefficient in the air phase	20,000	cm ² day ⁻¹
<i>λ</i>	Dispersivity	10	cm
<i>P</i> ₅₀	Pressure head at which 50% reduction in water uptake occur	-500	cm
<i>P</i> ₃	Exponent in the root water uptake response function	3	–